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- (71) Applicant (for all designated States except US): METAPROTEOMICS, LLC [US/US]; 9770 44th Avenue, N.W., Suite 100, Gig Harbor, WA 9833 (US).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): TRIPP, Matthew, L. [US/US]; 4202 71st Avenue Court NW, Gig Harbor, WA 98335 (US). BABISH, John, G. [US/US]; 508 White Church Road, Brooktondale, NY 14817 (US). BLAND, Jeffrey [US/US]; P.O. Box 477, Fox Island, WA 98333 (US). HALL, Amy, Jennae [US/US]; 11325 17th Avenue Court, NW, Gig Harbor, WA 98332 (US). KONDA, Veera [IN/US]; 5325 64th Avenue N.W., Gig Harbor, WA 98335 (US). PACIORETTY, Linda [US/US]; 508 White Church Road, Brooktondale, NY 14817 (US). DESAI, 98335 (US).

- (74) Agents: KUSMER, Toby, H. et al.; McDermott Will & Emery LLP, 28 State Street, Boston, MA 02109 (US).
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(54) Title: PROTEIN KINASE MODULATION BY HOPS AND ACACIA PRODUCTS

(57) Abstract: Botanical compounds to modulate kinase activity are disclosed. The compounds and methods disclosed also inhibit expression of COX-2, inhibit synthesis of prostaglandins selectively in target cells, and inhibit inflammatory response selectively. The compositions contain at least one fraction isolated or derived from hops or Acacia.

INTERNATIONAL SEARCH REPORT

International application No. PCT/US 06/47198

| A. CLASSIFICATION OF SUBJECT MATTER IPC(8) - A61K 38/43; A61K 36/00 (2007.10) USPC - 424/94.1; 424/725; 424/778 According to International Patent Classification (IPC) or to both national classification and IPC | | | | | | | |
|---|---|--|---|--|--|--|--|
| | | | | | | | |
| B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC(8) - A61K 38/43 (2007.10) USPC - 424/94.1, 424/725. 424/778 | | | | | | | |
| Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched IPC(8) - A61K 38/43; A61K 36/00 (2007.10) USPC - 424/94.1, 725, 778, 779, 775 | | | | | | | |
| Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) PubWest, DialogPRO, Google Patent, Google Scholar, PubMed/Medline, WIPO Search Terms Used: , kinase, kinase activity, hops, Acacia, cox\$, modulate, and combinations thereof. | | | | | | | |
| C. DOCU | MENTS CONSIDERED TO BE RELEVANT | | | | | | |
| Category* | Citation of document, with indication, where app | propriate, of the relevant passages | Relevant to claim No. | | | | |
| X | US 2003/0180402 A1 (Jia et al.) 25 September 2003 (2 para [0014], [0023], [0032], [0062], [0072], [0078]. | 5.09.2003). Entirety. Esp., abstract, | 1-2, 10-13, 17-18, 26-29 3-9, 14-16, 19-25,30-34 | | | | |
| Y | US 2005/0129791 A1 (Babish et al.) 16 June 2005 (16.0 [0060], [0119], [0120]. | 06.2005). Esp., abstract, para [0019], | 8,15,24,31,33-34 | | | | |
| Y | US 2004/0115290 A1 (Tripp et al.) 17 June 2004 (17.06 [0062], [0063], [0066]. | 4-7, 20-23 | | | | | |
| Y | US 2005/0192356 A1 (Babish et al.) 1 September 2005 [0016], [0017], [0080]. | 16,32 | | | | | |
| Y | US 2005/0042317 A1 (Babish et al.) 24 February 2005 [0040]. | 14,30 | | | | | |
| Y | Ward et al., "Therapeutic Potential of Phosphoinositide Biology (March 2003) vol. 10, pages 207-213. Esp. pages 207-213. | 3-Kinase inhibitors*, Chemistry & ge 209, para 3; page 210, para 1. | 3,19 | | | | |
| Y | Stevens et al., "Xanthohumol and Related Prenylflavon health", Phytochemistry (May 2004) vol. 65, page 131 | 9,25 | | | | | |
| | | | | | | | |
| Furth | er documents are listed in the continuation of Box C. | | | | | | |
| "A" docum | categories of cited documents: ent defining the general state of the art which is not considered | "T" later document published after the interdate and not in conflict with the applications are the properties. | Brigh but cited to underseand | | | | |
| to be o "E" earlier filing o | f particular relevance application or patent but published on or after the international late. | the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive | | | | | |
| cited to special | ent which may throw doubts on priority claim(s) or which is constablish the publication date of another citation or other reason (as specified) ent referring to an oral disclosure, use, exhibition or other | considered to involve an inventive step when the document is combined with one or more other such documents, such combination | | | | | |
| means "P" docum | ent published prior to the international filing date but later than | being obvious to a person skilled in th | е вл | | | | |
| Date of the actual completion of the international search Date of mailing of the international search report | | | | | | | |
| | 26 November 2007 (26.1.2007) | 20 DEC 2007 | | | | | |
| | nailing address of the ISA/US | Authorized officer: | | | | | |
| Mail Stop PC | CT, Attn: ISA/US, Commissioner for Patents 50, Alexandria, Virginia 22313-1450 | Lee W. Young | | | | | |
| | Vo. 571-273-3201 | PCT Helpdesk: 571-272-4300 PCT OSP: 571-272-7774 | | | | | |

PATENT COOPERATION TREATY

| From the INTERNATIONAL SEARCHING AUTHORITY | | | | | | | |
|--|-------------------------------------|---|--|--|--|--|--|
| To: Toby H. Kusmer McDermott Will & Emery LLP 28 State Street | | PCT WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY | | | | | |
| Boston, MA 02109 | | | | | | | |
| | | (PCT Rule 43bis.1) | | | | | |
| | Date of mailing (day/month/year) | 2 0 DEC 2007 | | | | | |
| Applicant's or agent's file reference 068911-0173 | | FOR FURTHER ACTION See paragraph 2 below | | | | | |
| International application No. International filing d | ate (day/month/year) | Priority date (day/month/year) | | | | | |
| PCT/US 06/47196 11 December 20 | 06 (11.12.2006) | 09 December 2005 (09.12.2005) | | | | | |
| International Patent Classification (IPC) or both national classification and IPC IPC(8) - A61K 38/43, 36/00 (2007.10) USPC - 424/94.1; 424/725; 424/778 | | | | | | | |
| Applicant Metaproteomics, LLC | | ! | | | | | |
| | | | | | | | |
| 1. This opinion contains indications relating to the following items: Box No. Basis of the opinion | | | | | | | |
| If a demand for international preliminary examination is made, this opinion will be considered to be a written opinion of the International Preliminary Examining Authority ("IPEA") except that this does not apply where the applicant chooses an Authority other than this one to be the IPEA and the chosen IPEA has notified the International Bureau under Rule 66.1 bis(b) that written opinions of this International Searching Authority will not be so considered. | | | | | | | |
| If this opinion is, as provided above, considered to be a written opinion of the IPEA, the applicant is invited to submit to the IPEA a written reply together, where appropriate, with amendments, before the expiration of 3 months from the date of mailing of Form PCT/ISA/220 cr before the expiration of 22 months from the priority date, whichever expires later. For further options, see Form PCT/ISA/220. | | | | | | | |
| 3. For further details, see notes to Form PCT/ISA/220. | | | | | | | |
| Name and mailing address of the ISA/US Date of completion Mail Stop PCT, Attn: ISA/US Commissioner for Patents 26 November 2 | of this opinion 2007(26.11.2007) | Authorized officer: Lee W. Young | | | | | |
| P.O. Box 1450, Alexandria, Virginia 22313-1450 Facsimile No. 571-273-3201 | | PCT Helpdesk: 571-272-4300 PCT OSP: 571-272-7774 | | | | | |

Form PCT/ISA/237 (cover sheet) (April 2007)

PCT/US2006/047196 20.12.2007

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No.

PCT/US 06/47196

| Box | No. I | Basis of this opinion |
|-----|---------|--|
| 1. | With re | egard to the language, this opinion has been established on the basis of: |
| | X | the international application in the language in which it was filed. |
| | | a translation of the international application into which is the language of a translation furnished for the purposes of international search (Rules 12.3(a) and 23.1(b)). |
| 2. | | This opinion has been established taking into account the rectification of an obvious mistake authorized by or notified to this Authority under Rule 91 (Rule 43bis.1(a)) |
| 3. | | egard to any nucleotide and/or amino acid sequence disclosed in the international application, this opinion has been shed on the basis of: |
| | a. typ | e of material |
| | | a sequence listing |
| | | table(s) related to the sequence listing |
| | b. for | mat of material |
| | J. 101 | on paper |
| | F | in electronic form |
| | | |
| | c. tin | ne of filing/furnishing |
| | | contained in the international application as filed . |
| | | filed together with the international application in electronic form |
| | | furnished subsequently to this Authority for the purposes of search |
| 4. | | In addition, in the case that more than one version or copy of a sequence listing and/or table(s) relating thereto has been filed or furnished, the required statements that the information in the subsequent or additional copies is identical to that in the application as filed or does not go beyond the application as filed, as appropriate, were furnished. |
| 5. | Addir | ional comments: |
| J. | Auuli | iona, aciminana. |
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PCT/US2006/047196 20.12.2007

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No.

PCT/US 06/47196

| Box No. V Reasoned statement un citations and explanati | | bls.1(a)(i) with regard to novelty, inventive step or industrial appling such statement | icability; |
|---|---|--|--|
| I. Statement | | | |
| Novelty (N) | Claims | 3-9, 14-16, 19-25, 30-34 | YES |
| 140VCity (14) | Claims | 1-2, 10-13, 17-18, and 26-29 | _ NO |
| | • | none | VEC |
| Inventive step (IS) | Claims Claims | 1-34 | YES NO |
| | | | |
| Industrial applicability (IA) | Claims | 1-34 | YES |
| | Claims | none | NO |
| (hereinafter 'JIA'). Regarding claims 1 and 17, JIA describes modulating the activity of a plurality of dis in need thereof, wherein said protein kina | a method (pa ease associat se modulation subject in ne | r PCT Article 33(2) as being anticipated by US 2003/0180402 A1 to JI. ara [0023]) and a composition (para [0033], [0035], [0056]), respective ted protein kinases (abstract COX-2 mediated diseases, para [0031] in is beneficial to the health of the subject (abstract; para [0031]); said and a therapeutically effective amount of a composition comprising a composition comprising a composition. | ely, for]) in a subject method |
| Regarding claims 2 and 18, JiA teaches t [0014]). | he method an | nd composition of claims 1 and 17, respectively, for inflammatory disor | ders (para |
| Regarding claims 10 and 26, JIA teaches is derived from Acacla nilotica (para [001- | | and composition of claims 1 and 17, respectively, wherein the compou | nd or extract |
| Regarding claims 11 and 27, JIA teaches compound is from Acacia nilotica extract | | and composition of claims 1 and 17, respectively, wherein the Acacla i | nilotica |
| Regarding claims 12 and 28, JIA teaches Acada nilotica extract is from acidified water acetate (para [0078]). | the method a ster(acidic), ed | and composition of claims 1 and 17, respectively, wherein the Acacia queous(polar) extractions (para [0062]), and organic extractions such | catechu or as and ethyl |
| Regarding claims 13 and 29, JIA teaches acceptable excipients are employed that | the method a can be agents | and composition of claims 1 and 17, respectively, wherein pharmacolos of color or absorption (para [0072]). | gically |
| Claims 16 and 32 lack an inventive step al. (hereinafter .BABISH'356.). | under PCT Art | ticle 33(3) as being obvious over JIA, in view of US 2005/0192356 A1 | to Babish et |
| teaches a composition comprising extractional specification and prostagiand cycloxygenase enzymes and prostagiand teach the use of acacia extracts, it was known to be acacia nilotica heartwood powder extracts. | ts isolated from 3:1 (para [00] lin synthesis a nown that extra BABISH'356, velop the met . One would l | A teaches as given above for claims 1 and 17, respectively. BABISH'm a natural plant (hops) wherein two different extracts (rho-isoalpha a 80]). These compounds exhibit anti-inflammatory action (abstract) inflammatory processes (para [0016], [0017]). Although BABISH racts of acada also exhibit anti-inflammatory action, as taught by JIA (it would have been obvious to one of ordinary skill in the art through thod of claim 18 and composition of claim 32 comprising a 5:1 ratio of have been motivated to do so to develop a more effective method of the east based on the teachings of JIA and BABISH'356. | icid, RIAA; and luencing I.358 does not (para [0014]). standard RIAA to |
| Claims 8, 15, 24, 31, 33, and 34 lack an i to Babish et al. (hereinafter 'BABISH'791' | nventive step). | under PCT Article 33(3) as being obvious over JIA, in view of US 200 |)5/0129791 A1 |
| Regarding claims 8 and 24, refer to the te the use of xanthohumol (para[0019]) in a | eachings of Jla formulation to | A as given above for claims 1 and 17, respectively. BABISH.791 furth provide anti-inflammatory effects (abstract). | ner teaches |
| the use of alpha and beta acids (para [00 treatment of disorders such as diabetes (| 19]), as given para (0060}). I in the art to c a more effect | JIA as given above for claims 1 and 17, respectively. BABISH.791 fur above in claims 1 and 17, having anti-inflammatory effects (abstract Based on the teachings of JIA, in view of the teachings of BABISH79 develop a method and composition comprising an anti-diabetic drug. live synergistic composition for treatment and would have had a reason. |) in the 91, it would One would |
| <u></u> | s | see continuatiuon sheet——————————————————————————————————— | |